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- GMELINS HANDBUCH DER ANORGANIS-CHEN CHEMIE, Auflage 8, 1927, pages 234-235, Verlag Chemie, Berlin, DE; "Chlor-System Nummer 6" z. Electroch. 13 (1907) 437
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Description

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The present invention relates to the production of chlorine dioxide by the electrolysis of highly acidic sodium chlorate solutions.

Chlorine dioxide is used as a bleach in a variety of environments, notably in the bleaching of wood pulp. Various chemical processes for the generation of chlorine dioxide by reduction of sodium chlorate in aqueous acid media have been described in the past and are in commercial operation. The chemical process may be depicted by the equation (I):

 $2ClO_3^- + 2Cl^- + 4H^+ \rightarrow 2ClO_2 + Cl_2 + 2H_2O$ (I)

U.S. Patents Nos. 4,426,263 and 4,362,707 (Hardee et al) describe an electrolytic process for producing chlorine dioxide using an electrocatalyst comprising a platinum group metal oxide as a cathode coating in an electrolytic cell containing sodium chlorate and sulphuric acid. This patent also describes the use of platinum group metal oxides as a catalyst in the absence of applied electrical current and indicates that this latter procedure is the preferred embodiment.

Since the material used as the cathode coating is itself a catalyst for the production of chlorine dioxide, the effect of an applied current cannot readily be determined but the data presented in the patent and also in an article by Hardee describing the electrochemical process of the patent (see "The Electrochemical Generation of Chlorine Dioxide Utilizing Electrolytic Oxide Coatings", Extended Abstracts, vol. 85-1, pp.617 to 618, The Electrochemical Society, 1985) suggest little beneficial effect of the applied current on the generation of chlorine dioxide.

According to the article, the better efficiencies are observed at lower current values and hence at lower contributions of electrolysis to the overall process of generation of chlorine dioxide. In particular, current efficiencies as low as 20% were observed at higher current densities. The observed loss in efficiency was believed to arise from further reduction of chlorine dioxide.

Poor results obtained in electrolytic experiments carried out at higher current densities are in a good correspondence with the cyclic voltammograms reported in the above-noted Hardee article, where the maximum current density observed for the electroreduction of 0.5M NaClO₆ is less than 10 mA/cm², which is, by an order of magnitude, lower than expected for such a high concentration of reducible species.

The experimental data in the Hardee article indicates to one skilled in the art that this prior art process is not limited by the electrochemical step involving chlorate but rather by a chemical step in which an electroactive species, different from chlorate, is formed, which undergoes subsequently an electroreduction. Accordingly, the rate of reduction of chlorate ion to chlorine dioxide is limited by a chemical reaction rather than an electrochemical one and this chemical reaction can be accelerated by the presence of a catalyst, as described in the Hardee patents and article. Although the platinum metal oxide catalyst appears to enhance the rate of chemical conversion of chlorate ion to chlorine dioxide, its electrocatalytic properties have a detrimental effect on the electrochemical stability of the desired product, namely chlorine dioxide, when practical current densities are applied to electrodes having a surface of such platinum metal oxides.

The Hardee article also states that the platinum group metal oxides are the only materials which show activity for the reduction of chlorate and data is presented showing the alleged ineffectiveness of platinum.

In accordance with the present invention, there is provided an electrochemical process for the production of chlorine dioxide which is based on an autocatalytic cycle utilizing part of the product, namely chlorine dioxide, for generation of the next portion of the same product.

It has been surprisingly found that pure chlorine dioxide, without any substantial contamination by chlorine, can be produced by passing a cathodic current through an aqueous acid solution of chlorate ions having a total acidity greater than that of about 7 normal sulphuric acid and removing generated chlorine dioxide from the aqueous acid solution.

In the process of the invention, this result is achieved by (a) using a cathode constructed of an electrochemically-active material which is also chemically inert and does not catalyse the chemical formation of chlorine dioxide from the aqueous chlorate solution, in the absence of any electric current therethrough, and (b) maintaining a dissolved concentration in the range of about 0.01 to about 15 grams per litre, of chlorine dioxide in the aqueous acid solution throughout the process.

The mechanism of generation of chlorine dioxide by the electrochemical process of the invention is believed to involve chemical reaction between chlorate ions and electrolytically-produced short-lived chlorite ions to form chlorine dioxide. Part of the chemically-produced chlorine dioxide is electrochemically reduced to form the chlorite ions, while the remainder is removed from the solution as product.

The reactions which are thought to occur may be depicted, as follows:

$$Clo_2 + e \rightarrow Clo_2^ Clo_2^- + Clo_3 + 2H^+ \rightarrow Clo_2 + Clo_2 (†) + H_2O$$
overall: $Clo_3^- + e + 2H^+ \rightarrow Clo_2 + H_2O$

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From these equations, it will be seen that the process can be considered autocatalytic, in that generated chlorine dioxide is used to produce the active species for reduction of chlorate ions. A residual concentration of chlorine dioxide must be maintained in the aqueous acid solution to sustain the autocatalytic cycle.

If the cell as a whole is considered, then the anodic and cathodic reactions may be depicted as follows:

Cathode:
$$2Clo_3^- + 2e + 4H^+ \rightarrow 2Clo_2 + 2H_2O$$

Anode: H_2O - $2e$ + $\frac{1}{2}O_2$ + $2H^+$

Cell: $2Clo_3^- + 2H^+ \rightarrow 2Clo_2 + \frac{1}{2}O_2 + H_2O$ (II)

As may be seen from equation (II) in comparison with equation (I), the electrochemical process of the invention produces the same amount of chlorine dioxide while half the amount of water is produced and half the amount of acid is consumed, as compared to the chemical process. The chlorine dioxide which is produced in the process of the invention generally is substantially pure since the reactions involved do not produce chlorine.

No chlorine dioxide is produced from the acidic aqueous chlorate solution while it is in the cell in contact with the electrodes in the absence of an applied cathodic current. The cathode which is used in the process of the invention may be constructed of any convenient electro-conductive material which is chemically inert (i.e. has no catalytic properties) to the chemical production of chlorine dioxide by reduction of chlorate ions in the acid aqueous reaction medium, in contrast to the materials described in U.S. Patent Nos. 4,426,263 and 4,362,707 referred to above. Suitable cathode materials include the platinum group metals and, preferably in view of its cheapness and ease of use, carbon in any form, for example, graphite and vitreous carbon. The use of a carbon cathode also is advantageous, since it stabilizes the intermediate state, that is, the chlorite ions, against further electroreduction to a lower valency state, such as CIO⁻ or CI⁻.

As noted above, the chlorine dioxide which is produced electrochemically in this invention generally is obtained free from chlorine, since chlorine is not produced by the reactions depicted by the equations given above. Chlorine generation is possible only if chloride ions are present in the reaction medium.

The possibility exists for the production of chloride ions by the acidic decomposition of chlorite ions if the excess of chlorate ion in the acid medium is insufficient, in accordance with the following equation:

The chloride ion produced in this way then can react chemically with the chlorate ions in accordance with the reaction depicted in equation (I) above to produce chlorine as well as chlorine dioxide. Although chlorine dioxide still is formed, the coproduction of chlorine represents an inefficiency with respect to the production of chlorine dioxide from chlorate ions and also is a source of current inefficiency.

The electrochemical process of the invention may be carried out under a wide range of process conditions. Essential to the present invention is the provision of an aqueous acid electrolyte solution containing dissolved chlorate ions and having a total acidity greater than that of about 7 normal sulphuric acid. At acidities corresponding to below about 7 normal sulphuric acid, the production of pure chlorine dioxide is not possible.

The acidity may be provided most conveniently by sulphuric acid although any other strong mineral acid, other than hydrochloric acid, or a mixture of acids, may be employed, such as perchloric acid (HClO₄), or nitric acid (HNO₃).

Hydrochloric acid is avoided, since the introduction of chloride ions would produce the undesired side chemical reaction with chlorate ions to produce chlorine dioxide and chlorine. An acid aqueous chlorate solution having a total acidity corresponding to that of about 9 to about 11 normal sulphuric acid is preferred.

The chlorate ions in the electrolyte are provided preferably by sodium chlorate, since this chemical is the most readily-available form of chlorate. However, other alkali metal chlorates, such as potassium chlorate, lithium chlorate, rubidium chlorate and cesium chlorate may be used, as well as alkaline earth metal chlorates, such as beryllium chlorate, magnesium chlorate, calcium chlorate, strontium chlorate, barium chlorate and

radium chlorate, and mixtures of two or more of such chlorates. The concentration of chlorate ions in the electrolyte may vary widely from about 0.001 to about 7 molar, preferably about 0.1 to about 2 molar.

In order to sustain the reactions which are thought to be involved in the electrochemical process of the invention, it is essential to maintain a dissolved concentration of chlorine dioxide in the electrolyte throughout the process. Chlorine dioxide generation ceases if all the produced chlorine dioxide is removed. In addition, some dissolved chlorine dioxide is necessary at start up. A concentration of dissolved chlorine dioxide in the range of about 0.01 to about 15 grams per litre (gpl) may be employed, preferably about 0.1 to about 8 gpl, at the initial startup and during the reaction.

Although some inefficiency may be tolerated, in order to minimize side reactions which result in chlorine production and to maximize the overall chemical efficiency of the production of one mole of chlorine dioxide for each mole of chlorate ion consumed, the concentration of chlorate ion in the electrolyte should be in substantial excess to the concentration of dissolved chlorine dioxide, generally a molar excess of at least about 2:1, preferably at least about 10:1, usually up to about 1000:1.

Generally, the concentration of dissolved chlorine dioxide is maintained at a substantially uniform level during the process by removing chlorine dioxide at the rate of its formation. Chlorine dioxide spontaneously decomposes at high partial pressures thereof and it is necessary to dilute the chlorine dioxide well below the decomposition partial pressure, usually below about 100 mmHg (about 1.3 x 10⁴ Nm⁻²). Any convenient diluent gas, usually air, may be used to trip the generated chlorine dioxide from the electrolytic cell and to provide the required dilution. Chlorine dioxide may be recovered from the off-gas stream by dissolution in water.

The electrical potential applied to the cathode during the electrochemical reaction depends on the material of construction of the electrode and usually varies from about +1.0 to about -0.5 Volts as compared with a saturated calomel electrode (SCE). For a carbon electrode, the preferred potential is approximately +0.4 Volts while for a platinum electrode, the preferred potential is approximately +0.7 Volts. The process usually is operated under constant voltage conditions while the current also preferably is constant.

The temperature of operation of the cell affects the purity of the chlorine dioxide gas which is obtained. Higher temperatures favour the formation of chloride ions by decomposition of chlorite ions, as described above, in accordance with the equation:

As discussed above, formation of chloride lons in this way results in the formation of chlorine, with the consequent loss of efficiency and chlorine dioxide purity. Accordingly, it is preferred to operate at temperatures of about 20° to about 25°C.

The process of the present invention may be carried out in any convenient cell arrangement in which anode and cathode electrodes are located and between which current may be passed. The cell may be divided physically into anolyte and catholyte chambers by any convenient cation-exchange membrane. With a divided cell arrangement, the aqueous acid chlorate solution is fed to the cathode compartment while water is fed to the anode compartment, the latter containing an electrolyte, such as an acid solution.

However, operation in an undivided cell or a cell with a simple non-membrane separator also is possible. As noted above, the electrochemical reaction at the cathode surface is believed to be the formation of chlorite ion from chlorine dioxide. In an undivided cell, such chlorite ions will attempt to migrate to the anode but are consumed by the chlorate ions present in large excess in the electrolyte to form chlorine dioxide, so that the chlorite ions should never reach the anode and, in addition, their lifetime in the acidic medium is very short.

The generation of chlorine dioxide by the process of the invention is accompanied by the formation of by-products. As noted earlier, the anodic reaction in the cell produces gaseous oxygen, which may be vented in any convenient manner. The other by-products are water produced by the electrochemical reaction and a salt of the cation of the chlorate and the anion of the acid consumed in the process. These may be removed respectively by any convenient procedure, such as by evaporation and crystallization outside the cell.

Chlorine dioxide is produced in pure form from an aqueous acid chlorate solution by passing a cathodic current through the solution from a cathode constructed of material chemically inert with respect to the formation of chlorine dioxide from the solution. Such a result is achieved by maintaining a dissolved concentration of chlorine dioxide in the solution.

In the Examples which follow, reference is made to the accompanying drawing, in which:

Figure 1 contains a series of three voltammetric curves obtained in experiments described in these Examples.

The invention is illustrated by the following Examples:

Example 1

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Cyclic voltammetric studies were effected on an aqueous solution containing about 0.1 g/L of chlorine

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dioxide, 1 M NaClO₃ and 10 N H₂SO₄, using, in one case, a glassy carbon electrode and, in another case, a platinum disc electrode, each having a surface area of 0.196 cm². The current was plotted against the applied potential and the results are reproduced as curves a (glassy carbon) and b (platinum) in Figure 1. The initial potential applied was +1.0V vs. SCE and a sweep rate of 0.1 Vs⁻¹ was used. A further run was made using glassy carbon on a solution from which the sodium chlorate was absent. These results are reproduced as curve c in Figure 1.

It will be seen from the data presented in curves a and b in Figure 1 that both the ClO₂ reduction current to ClO₂⁻, which is proportional to the ClO₂ concentration, and corresponding reoxidation current of ClO₂⁻ back to ClO₂, recorded during consecutive potential scans between +1.0V and +0.6V vs. SCE, increases substantially with the duration of the multicyclic experiments. This result indicates a progressive accumulation or self-perpetuated multiplication of chlorine dioxide in the proximity of the electrode, both for the glassy carbon and platinum electrode. In comparison, curve c of Figure 1 shows no accumulation of chlorine dioxide in the absence of the chlorate ions.

Example 2

Electrolytic studies were carried out in a divided H-cell using reticulated vitreous carbon foam as the cathode material and platinum foil as the anode material. A potentiostatic mode of operation was adopted at a cathodic potential of +0.2 volt vs. Hg/Hg₂SO₄ as the reference electrode.

The catholyte of volume approximately 100 ml contained about 10N H₂SO₄, about 1M NaClO₃ and a variable initial dissolved ClO₂ concentration. During electrolysis, gaseous products, ClO₂ and Cl₂ were stripped to a potassium iodide (KI) trap by bubbling nitrogen and by applying a low level of vacuum. Both the nitrogen flow and the vacuum were adjusted to maintain a substantially constant level of electrolyte in the compartments and, at the same time, to maintain a substantially constant concentration of dissolved chlorine dioxide in the catholyte, so that the stripping rate of chlorine dioxide was approximately equal to the production rate of chlorine dioxide.

The electrolyte was analyzed for CiO₂, Cl₂, ClO₃, Cl, CiO₂ and acidity both before and after electrolysis while the KI trap was analyzed and changed every 30 to 60 minutes.

The results obtained are set forth in the following Table I:

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TABE I

AUTOCATALYTIC ELECTRONEDUCTION OF CIO, to CIO,

NO.	NO. INITIAL man	TAL AM	L AMOUNTS IN	NI	INITIAL NORMALITY	TEMP.	AVERAGE	CHARGE [C]	NCE	CUR	CHEMICAL EFFICIENCY	CIO2/CI2
	C10 ²	C1 ₂	C102 C12 NaC103 NaC1	NaC1	of H ₂ S0 ₄	[၁.]	[A]		% C]	ઝ્ લ	> €	RATTO
-	1.56	0.08	8.66	9.0	9.76	23	0,40	3000	86	88	83	9
2	1.56	0.08	8.66	9.0	9.76	श्च	0.43	4018	86	g	83	iğ.
က	2.91	0.19	97.9	9.0	9.86	Ю	0.58	3643	8	88	88	33
4	3.51	0.11	182.2	0.4	9.58	Ю	0.58	6166	ま	83	87	Ŕ
2	1.53	0.08	6.66	0.3	3.62	rs R	0.61	2669	103	83	83	Ŕ
9	1.58	0.10	99.5	0.4	86.6	88	979	11000	81	\$	%	12
7	1.33	D.19	9.101	9.0	5. 6	જી	92.1	10000	88		88	. 81

As may be seen from Table I, the chemical and current efficiencies which were obtained were very high, particularly at ambient temperature. The current required was dependent on both dissolved chlorine dioxide concentration and temperature, with the temperature effect being much more significant. As may be seen from run No. 4, a significant increase in sodium chlorate concentration did not significantly affect the process.

The mole ratio of ClO₂/Cl₂, and hence the purity of chlorine dioxide, although good in all the experiments, was much better at ambient temperature and is in a good correspondence with the temperature dependence of current efficiency.

Chlorite ions postulated as short-lived intermediates in the autocatalytic process were not detected either in the catholyte and analyte, before and after electrolysis.

Maintaining a residual chlorine dioxide concentration was critical for the operation of the process. In an experiment where all the chlorine dioxide was stripped from the solution, no further electrogeneration of chlorine dioxide was observed and the current measured under potentiostatic conditions went to zero.

In summary of this disclosure, the present invention provides a novel method of producing chlorine dioxide by an autocatalytic cathodic electrochemical reduction of chlorate ions.

Claims

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- 1. An electrochemical process for the production of chlorine dioxide by passing a cathodic electrical current through an aqueous acid solution of chlorate ions having a total acidity greater than that of about 7 normal sulfuric acid and removing generated chlorine dioxide from the aqueous acid solution, characterized by:
 - (a) using a cathode constructed of an electrochemically-active material which is also chemically inert and does not catalyse the chemical formation of chlorine dioxide from the acidic chlorate solution in the absence of any electric current therethrough, and
 - (b) maintaining a dissolved concentration, in the range of about 0.01 to about 15 grams per litre, of chlorine dioxide in said aqueous acid solution throughout said process.
- 2. The process claimed in claim 1, characterized by maintaining a molar excess concentration of chlorate ions with respect to the concentration of dissolved chlorine dioxide in said aqueous acid solution of at least about 2:1.
- 3. The process claimed in claim 2, characterized in that the molar excess concentration is up to about 1000:1.
- 4. The process claimed in claim 2 or 3, characterized in that molar excess concentration is at least about 10:1, the aqueous acid solution has a chlorate concentration of about 0.1 to about 2 molar and said aqueous acid solution contains about 9 to about 11 normal sulphuric acid.
- 5. The process claimed in any of claim 1 to 4, characterized in that the concentration of dissolved chlorine dioxide is in the range of about 0.1 about 8 gpl (g/l).
- 6. The process claimed in any one of claims 1 to 5, characterized in that said dissolved concentration of chlorine is maintained substantially uniform in said aqueous acid solution throughout said process by removing generated chlorine dioxide from the aqueous acid solution at a rate corresponding to its rate of formation.
- 7. The process claimed in any one of claims 1 to 6, characterized in that said cathode is constructed of carbon.
- 8. The process claimed in any one of claims 1 to 7, characterized in that an electrode potential of about +1.0 to about -0.5 Volt as compared to a saturated calomel electrode is applied to the cathode.
- 9. The process claimed in any one of claims 1 to 8, characterized by being effected at a temperature less than about 40°C.

Patentansprüche

- 1. Electrochemisches Verfahren zur Herstellung von Chlordioxid, indem ein kathodischer elektrischer Strom durch eine wässrige saure Lösung von Clorationen geleitet wird, die eine gesamte Azidität aufweist, die höher ist als die von etwa 7 normaler Schwefelsäure, und indem das erzeugte Chlordioxid aus der sauren wässrigen Lösung entfernt wird, dadurch gekennzeichnet, daß
 - (a) eine Kathode aus einem elektrochemisch aktiven Material verwendet wird, das außerdem chemisch inert ist und die chemische Bildung von Chlordioxid aus der sauren Chloratiösung in Abwesenheit eines durchgehenden elektrischen Stromes nicht katalysiert; und
 - (b) die gelöste Konzentration des Chlordioxids in der wässrigen sauren Lösung während des ganzen Verfahrene im Bereich von etwa 0,01 bis etwa 15 g pro Liter gehalten wird.

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- 2. Verfahren nach Anspruch 1, dadurch gekennzeichnet, daß eine molare Überschußkonzentration von Chlorationen in Bezug auf die Konzentration des in der sauren wässrigen Lösung gelösten Chlordioxids von mindestens etwa 2:1 aufrechterhalten wird.
- 3. Verfahren nach Anspruch 2, dadurch gekennzeichnet, daß die molare Überschußkonzentration bis zu etwa 1000;1 beträgt.
- 4. Verfahren nach einem der Ansprüche 2 und 3, dadurch gekennzeichnet, daß die molare Überschuß-konzentration mindestens etwa 10:1 beträgt, wobei die wässrige saure Lösung eine Chloratkonzentration von etwa 0,1 bis etwa 2 molar aufweist und die wässrige saure Lösung etwa 9 bis etwa 11 normale Schwefölsäure enthalt.
- 5. Verfahren nach einem der Ansprüche 1 bis 4, dadurch gekennzeichnet, daß die Konzentration des gelösten Chlordioxide im Bereich von etwa 0,1 bis etwa 8 g/l liegt.
- 6. Verfahren nach einem der Ansprüche 1 bis 5, dadurch gekennzeichnet, daß die gelöste Konzentration des Chlordioxids in der wässrigen sauren Lösung während des ganzen Verfahrens im wesentlichen konstant wird, indem das gebildete Chlordioxid aus der wässrigen sauren Lösung mit einer Geschwindigkeit entfernt wird, die seiner Bildungsgeschwindigkeit entspricht.
- 7. Verfahren nach einem der Ansprüche 1 bis 6, dadurch gekennzeichnet, daß daß die Kathode aus Kohlenstoff besteht.
- 8. Verfahren nach einem der Ansprüche 1 bis 7, dadurch gekennzeichnet, daß an die Kathode ein Elektrodenpotential von etwa +1,0 bis etwa -0,5 Volt in Bezug auf eine gesättigle Kalomelektrode angelegt wird.
- 9. Verfahren nach einem der Ansprüche 1 bis 8, dadurch gekennzeichnet, daß es bei einer Temperatur von weniger als etwa 40°C durchgeführt wird.

Revendications

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- 1. Procédé électrochimique pour la production de dioxyde de chlore en faisant passer un courant électrique cathodique dans une solution aqueuse acide d'ions chlorate ayant une acidité totale supérieure à celle de l'acide sulfurique environ 7 N et en éliminant le dioxyde de chlore produit de la solution aqueuse acide, caractérisé en ce que:
 - (a) on utilise une cathode constituée d'un matériau électrochimiquement actif qui est également chimiquement linerte et qui ne catalyse pas la formation chimique de dioxyde de chlore au départ de la solution acide de chlorate en l'absence de tout courant électrique dans celle-ci, et
 - (b) on maintient une concentration de dioxyde de chlore dissous dans ladite solution aqueuse acide comprise dans la gamme d'environ 0,01 à environ 15 grammes par litre pendant tout ledit procédé.
- 2. Procédé selon la revendication 1, caractérisé en ce que l'on maintient une concentration molaire en excès d'ions chlorate par rapport à la concentration de dioxyde de chlore dissous dans ladite solution aqueuse acide, qui est d'au moins environ 2:1.
- 3. Procédé selon la revendication 2, caractérisé en ce que la concentration molaire en excès va jusqu'à environ 1000:1.
- 4. Procédé selon la revendication 2 ou 3, caractérisé en ce que la concentration molaire en excès est d'au moins environ 10:1, que la solution aqueuse acide a une concentration en chlorate comprise entre environ 0,1 et environ 2 molaire et que ladite solution aqueuse acide contient de l'acide sulfurique depuis environ 9 N jusqu'à environ 11 N.
- 5. Procédé selon l'une quelconque des revendications 1 à 4, caractérisé en ce que la concentration du dioxyde de chlore dissous est comprise dans la gamme de 0,1 à environ 8 gpl (grammes par litre).
- 6. Procédé selon l'une quelconque des revendications 1 à 5, caractérisé en ce que ladite concentration de dioxyde de chlore dissous est maintenue essentiellement uniforme à ladite solution aqueuse acide pendant tout ledit procédé, en éliminant le dioxyde de chlore produit de la solution aqueuse acide à un débit correspondant à son débit de formation.
- 7. Procédé selon l'une quelconque des revendications 1 à 6, caractérisé en ce que ladite cathode est constituée de carbone.
- 8. Procédé selon l'une quelconque des revendications 1 à 7, caractérisé en ce qu'on applique à la cathode un potentiel d'électrode compris entre environ +1,0 et environ -0,5 V par rapport à une électrode saturée au calomel.
- 9. Procédé selon l'une quelconque des revendications 1 à 8, caractérisé en ce qu'on le met en oeuvre à une température inférieure à environ 40°C.

FIGURE 1.

